Infection of the lymphatic system by Aureobasidium pullulans in a patient with erythema nodosum leprosum

Authors

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ABSTRACT

Aureobasidium pullulans is a causal agent of phaeohyphomycosis, occasionally found in men and animals. As an agent of different opportunistic fungal processes, it may cause fungemia, systemic infections and abscesses in different viscera. This paper aims to report a case of a patient with infection of the lymphatic system by *A. pullulans*. A 23-year-old patient being treated for erythema nodosum leprosum presented a 60-day complaint of daily fever, hoarseness, odynophagia and weight loss. Laboratory tests showed pancytopenia with severe neutropenia, cervical adenomegaly and solid contrast uptake lesion in the oropharyngeal region. Due to neutropenia and sepsis the patient was initially treated with cefepime and vancomycin, but there was no clinical improvement. Lymph node puncture-aspiration showed yeast-form fungus identified as *A. pullulans* by sequencing ITS region. The patient was treated with amphotericin B deoxycholate, leading to complete recovery of bone marrow function and regression of adenomegaly and the oropharyngeal lesion.

Keywords: HIV; pneumonia; macrolides; mortality.

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INTRODUCTION

Aureobasidium pullulans is a fungal agent of universal distribution, an occasional causing agent of phaeohyphomycosis in human beings and animals. It is a black yeast-form fungus that causes many opportunistic fungal processes and owing to its capacity to produce extracellular biodegradable polysaccharide pullulan, as well as many different hydrolytic enzymes (e.g. amylase, proteinase, cellulase), it is considered an agent of great biotechnological importance.¹

A. pullulans is an opportunistic agent rarely associated with infections in human beings and its pathogenic mechanisms are not very well known. However, infections located in the most varied organic topographies have already been associated with the agent, among which there are keratomycosis, spleen and mandible abscesses, meningitis, skin mycosis and peritonitis.²⁻⁶

Systemic fungal infections are relatively common in patients receiving broad spectrum antibiotics, using invasive devices, as well as those immunosuppressed. Among febrile neutropenic patients, systemic fungal infection is identified in about 10% of non-responders to initial antibacterial therapy, and in these patients opportunistic fungi such as *Candida* spp. and *Asper*- *gilus* spp. are the isolated agents in 80% to 90% of the cases.⁸ However, rare cases of septicemia by *A. pullulans* have also been reported.^{5,9-12}

The purpose of the present study was to report a case of a patient who had lymphatic system lesions which were attributed to infection by *A. pullulans*.

CASE REPORT

Male, 23-year-old patient, coming from Brasília, were being followed up by the Dermatology Division, University Hospital of Brasilia. The patient had received treatment for lepromatous leprosy seven years ago, although due to the erythema nodosum leprosum, he was still taking thalidomide (200 mg/day) and prednisone (5 mg/day) to achieve clinical control. For a period of 60 days the patient had been presenting symptoms of daily fever, hoarseness, odynophagia and weight loss which lead to the diagnosis of bacterial pharyngotonsillitis. The patient was initially treated as an outpatient having received prescription of amoxacilin, cephalexin and then clindamycin and ciprofloxacin, but with no symptoms relief.

At the hospital admission, the patient was markedly asthenic, febrile (39°C) and had

abnormality of vocal timbre. Upon clinical examination, he presented significant increase of volume and erythema of the palatine tonsils, as well as painful cervical adenomegaly. Clinical thoracic and abdominal examination were normal, except for dullness to percussion over Traube's space.

Laboratory tests showed a white blood cells count with pancytopenia and severe neutropenia, no findings in chest x-ray and no liver or kidney abnormalities. Thereafter, the detection of febrile neutropenia, in a context of sepsis with probable oropharyngeal focus, led to immediate introduction of antibiotic therapy with cefepime and vancomycin (2 g IV BID and 1 g IV BID, respectively). As the patient's health status kept on deteriorating even after 48 hours of antibiotics use, and considering that the patient came from a kala-azar endemic region, amphotericin B deoxycholate (IV, 50 mg/day) was associated to the treatment.

Further radiological investigation using CT scan showed a contrast-uptake solid lesion, involving the left side of the soft palate and palatoglossal and palatopharyngeal arches, measuring 30 mm in diameter, associated with multiple non-confluent adenomegaly of bilateral cervical chains (Figure 1). Abdominal CT scan confirmed the clinical findings, showing a mild increase in spleen volume. Myelogram showed hypocellular bone marrow with shift to the left, compatible with bone marrow aggression, without parasites or fungi in the analyzed sample. A rapid rK39 immunochromatographic dipstick test (Kala-azar Detect[™] Rapid Test, InBios International, Seattle, WA, USA) was also performed and turned out negative. Blood cultures did not grow fungal or bacterial agents and laryngoscopy revealed no findings suggestive of neoplasm.



Figure 1: CT scan with contrast of the neck. High-enhanced exophytic lesion in the soft palate region on the left. Presence of multiple enlarged lymph nodes of the neck chains (*).

Viral serology (hepatitis B and C, HIV) was negative.

Therefore, cervical lymph node fine needle aspiration was undertaken and the analysis of the collected material showed presence of many fungal yeast-form structures with sprouting that were phagocytized by reticular cells (Figure 2). At that time, Paracoccidioidis brasiliensis was ruled out by PCR. The culture of the same material revealed growth of yeast-form fungal colony, comprised by hyaline septated hyphae given rise synchronically to multiple conidia. Some attempts were made to define the etiological agent using molecular markers. PCR amplification and sequencing of intergene region ITS1/2 of gene 18S of the DNA extracted from the isolated colony led to the identification of A. pullulans as the putative pathogen. The fragment of sequenced ITS region (HUB_PAC1) presented 98% identification with the environmental strain of A. pullulans SN22 (GENBANK access number: FJ515165), as well as with the clinical strain UWFP 769 (GENBANK access number: AY213639), showing a high level of preservation of sequenced nucleotides when they were aligned (Figure 3). After 30 days of incubation at 35°C, the fungal colony started to produce a dark pigment (Figure 4) with presence of black septated hyphae, characteristic of A. pullulans.

The patient progressed with remarkable clinical recovery and had remission of fever and neutropenia after the 2^{nd} and 12^{th} day of amphotericin B use, respectively. After a total dose of 1,850 mg, administered in 40 days, the patient had already showed complete remission of adenomegaly and almost complete bone marrow recovery, persisting with mild anemia (Hb = 11.8g/dL). At that time, the patient was discharged using fluconazole at 300 mg, PO, twice a week for 8 weeks. Three months after the end of treatment, the patient was asymptomatic and had complete bone marrow recovery. Thalidomide was maintained during the whole period of treatment (100 mg/day).



Figure 2: Cytology of lymph node aspirate. Many yeast-like fungal forms of round and oval shape and varied sizes are seen. Some fungal structures demonstrate polar sprouting (Grocott; 400X).

	10	20	30	40	50	60	70	80	90	100
HUB_PAC1 A. pullulans_SN22 A. pullulans_UWFP769	GGTGCTCAGCGCCCG GGTGCTCAGCGCCCG GGGTGCTCAGCGCCCG	ACCTCCAACC ACCTCCAACC ACCTCCAACC	CTTTGTTGTT CTTTGTTGTT CTTTGTTGTT CTTTGTTGTT	AAAACTACCT AAAACTACCT AAAACTACCT	TGTTGCTTTGC TGTTGCTTTGC TGTTGCTTTGC	GCGGGACCCGC GCGGGACC-GC GCGGGACC-GC	TCGGTCTCGA TCGGTCTCGA TCGGTCTCGA	GCCGCTGGGG/ GCCGCTGGGG/ GCCGCTGGGG/	ATTOGTOCCA ATTOGTOCCA ATTOGTOCCA	LGGC LGGC LGGC
HUB_PAC1 A. pullulans_SN22 A. pullulans_UWFP769	110 GAGCGCCCGCCAGAGT GAGCGCCCGCCAGAGT GAGCGCCCGCCAGAGT	120 . TAA-CCAAAC TAAACCAAAC TAAACCAAAC	130 . TCTTGTTATT TCTTGTTATT TCTTGTTATT	140 . TAACCGGTCG TAACCGGTCG	150 . TCTGAGTTAAJ TCTGAGTTAAJ	160 . AATTTTGAATA AATTTTGAATA AATTTTGAATA	170 	180 	190 JATCTCTTGG JATCTCTTGG JATCTCTTGG	200 JTTC JTTC JTTC
HUB_PAC1 A. pullulans_SN22 A. pullulans_UWFP769	210 . TCGCATCGATGAAGAA TCGCATCGATGAAGAA TCGCATCGATGAAGAA	220 . CGCAGCGAAA CGCAGCGAAA CGCAGCGAAA	230 . TGCGATAAGT TGCGATAAGT	240 . AATGTGAATT AATGTGAATT AATGTGAATT	250 GCAGAATTCAG GCAGAATTCAG GCAGAATTCAG	260 TGAATCATCG TGAATCATCG TGAATCATCG	270 AATCTTTGAA AATCTTTGAA AATCTTTGAA	280 CGCACATTGCC CGCACATTGCC CGCACATTGCC	290 	300 ATT ATT ATT
HUB_PAC1 A. pullulans_SN22 A. pullulans_UWFP769	310 CCGAGGGGCATGCCTG CCGAGGGGCATGCCTG CCGAGGGGCATGCCTG	320 TTCGAGCGTC TTCGAGCGTC TTCGAGCGTC	330 ATTACACCAC ATTACACCAC ATTACACCAC	340 TCAAGCTATG TCAAGCTATG	350 CTTGGTATTGG CTTGGTATTGG CTTGGTATTGG	360 GEGCCGTCCT GEGCCGTCCT GETGCCGTCCT	370 TAGTTGGGCG TAGTTGGGCG TAGTTGGGCG	380 	390 ACCTCGGCGA ACCTCGGCGA	400 VGGC VGGC
HUB_PAC1 A. pullulans_SN22 A. pullulans_UWFP769	410 	420 . GTAGAT GTAGTAGAAT GTAGTAGAAT	430 . TTATTCGAAC TTATTCGAAC TTATTCGAAC	440 GTCTGTCAAA GTCTGTCAAA GTCTGTCAAA	450 	460 CTGCCGACTG CTGCCGACTG CTGCCGACTG	470 AAACCTTTTA AAACCTTTTA AAACCTTTTA AAACCTTT-A	480 	490 TGACCTCGG TGACCTCGG TGACCTCGG	500 JATC JATC JATC
HUB_PAC1 A. pullulans_SN22 A. pullulans_UWFP769	510 	520 TGAACT TGAACT TGAACT								

Figure 3: Alignment of sequencing of regions ITS 1/2 of gene 18S *A. pullulans*. Sequence HUB_PAC1, corresponds to studied strain, shows high level of similarity when compared to environmental strain SN22 (GENBANK access number: FJ515165) and strain of clinical origin UWFP769 (GENBANK access number: AY213639).



Figure 4: Macroscopy of fungal colony obtained in the material from lymph node aspirate. Yeast-like colonies of beige color, with initial points of dark pigment in areas after 30 days of incubation (agar-Sabouraud, 35°C).

DISCUSSION

A. pullulans is a saprophyte filamentous fungus of universal distribution, easily isolated in airborne samples, from superficial layers of the ground, domestic waste and leaves surface. It is a dematiaceous fungus, which presents colonies of quick growth, preferably at 25°C, capable of producing black pigment similar to melanin, which is deposited on the cell wall and may act as virulence factor inhibiting phagocytosis by the host's immune system.¹³ *A. pullulans* is normally considered a contaminant of biological samples, but the isolation from sterile sites should be valued given that the agent may act as an opportunistic agent.

When isolated in a culture, *A. pullulans* presents colonies initially yeast-form, yellowish, beige or light brown formed by dense groups of conidia synchronically originated from undifferentiated hyaline hyphae. As a result of maturation of colonies, hyphae accumulate pigment and become dark brown.^{7,11} The diagnosis is made by isolation and morphological characterization in specific culture medium which may be successfully studied, but their yeast-form shape may be mistaken by other yeasts of Dothideomycete class.^{2,6}

The constant advance of medical science has provided increased survival to severely ill patients who in some situations require aggressive treatments, which commonly lead to iatrogenic immunosuppression, the use of invasive intrabody devices, and the need to use broad spectrum antibacterial agents for prolonged periods of time. Nevertheless, all those conditions are recognized as risk factor for fungal infections and have contributed for the increase in frequency of infections by opportunistic fungal agents.¹⁴ There are seven reported cases of systemic disseminated infections by A. pullulans as a result of isolation of the agent in blood culture. Joshi et al.5 reported a case of a 11-year-old patient who had Fanconi anemia and, during the conditioning regimen for the performance of bone marrow transplantation, because of bone marrow suppression, developed a febrile episode followed by cutaneous lesions caused by A. pullulans, which was later isolated in blood culture collected from the venous central catheter. Another case was reported by Kaczmarski et al.,12 who described a 28-year-old patient with acute myeloid leukemia, who had neutropenia secondary to chemotherapy and reported isolation of A. pullulans in a blood sample collected from Hickman catheter used to infuse chemotherapy agents. The involvement of these neutropenic patients, as well as the large proportion of cases of peritonitis caused by dialysis catheter infection, confirm the role of invasive intrabody devices and immunosuppression states as facilitators of these systemic infections.15

In the initial assessment, the significant findings of patient were increased volume and erythema of palatine tonsils, cervical adenomegaly and splenomegaly, associated with hematological manifestation of pancytopenia. The severe and extensive involvement of the lymphohematopoietic system, together with absence of clinical improvement after 48 hours after onset of antimicrobials, prompted to the introduction of amphotericin B, in addition to broad the possibilities of differential diagnoses.

Clinical data supported by the patient coming from regions endemic of both kala-azar and South-American paracoccidioidomycosis and also the suspicion of hematological neoplasm (lymphoma) or squamous cell carcinoma, led to immediate myelogram and lymph node puncture-aspiration for diagnostic investigation.

Even though the myelogram had shown neither fungal nor parasite agents or neoplastic findings, mycological diagnosis was suggested through fine needle aspiration of a cervical lymph node, which showed yeast-like fungal structures with sprouting which were phagocytized by host macrophages. From a cytopathology standpoint, owing to absence of demaceous hyphae in the material, it was not possible to define an etiological classification due to the similarity of A. pullulans conidia and those of other yeast-like fungi.¹⁶ In the past, the production of pigment was a key factor for the identification of demaceous species such as Aureobasidium from culture medium, but it is known that this event occurs only as a result of hyphae aging, which delays or even causes diagnostic errors. In view of these factors, the use of gene sequencing in regions ITS1/2 showed quick and precise etiological diagnosis of A. pullulans, which reinforces the efficacy of this method as a diagnostic tool for fungal species, in agreement with the data reported in the literature.^{17,18}

Salkin et al.² reported a case of fungal invasion of the lymphatic system, causing the formation of spleen abscess in a patient systematically affected by histiocytic lymphoma. The entry of the infection was not identified by the authors, but the profound immunesuppression caused by the lymphoma seemed to have played a predisposing role in the secondary infection by A. pullulans. In another case, disseminated infection of the fungus was reported in a 28-year-old patient who had multiple exposed fractures of the left femur and tibia because of a car accident.¹¹ The exposure of the wounds to the ground during the car accident was reported by the authors as a likely source of infection, since the ground is a known reservoir of the agent. A retrospective analysis of our case made us believe that the lesions in the oral mucosa could have served as an entry point for the infection in our patient, since there were no other specific foci. Moreover, immune studies have shown that patients with polar forms of lepromatous leprosy and erythema nodosum leprosum present a pattern of immune response mediated by Th2 lymphocytes with the production of IL4 and IL10 cytokines. Such cytokines induce the humoral immune response via CD8(+) lymphocytes and also seem to lead to negative regulation of Toll-like 2/1 receptors, determining an ineffective response against Hansen bacilli.¹⁹ Both the history of lepromatous leprosy of the patient and the concurrent use of

thalidomide at the moment of infection, a drug known for its anti-TNF-alpha action, made us advocate that the patient might have presented a predominantly humoral immune response pattern to *A. pullulans*. These factors may have facilitated the dissemination of infection in view of low levels of Th1-type (INF gamma, TNF-alpha) proinflammatory cytokines, widely known for their important immune response against fungal agents.²⁰

Treatment of *A. pullulans* infections, either localized or disseminated, is not a consensus since up to the present day treatment recommendations are based on case reports and case series. In the present case, the total dose of 1,850 mg of conventional amphotericin B was enough for complete remission of cervical adenomegaly and bone marrow recovery. Nevertheless, the patient's prompt recovery after introduction of the antifungal, even before the isolation of the agent, reinforces the importance of following up evidence-based protocols for sepsis management, especially in the immunosuppressed or neutropenic hosts.

Therefore, even in the absence of fungal structures visualized in the analysis of bone marrow aspirate, we believe that the manifestation of pancytopenia presented by the patient may have been secondary to bone marrow invasion by the agent. To support this hypothesis, we excluded other infectious agents that may have determined similar clinical and laboratory manifestations. Alternatively, medications such as clindamycin,²¹ ciprofloxacin,²² and thalidomide²³ had been associated with episodes of bone marrow suppression, but considering the rarity of these reports and the complete hematological recovery of the patient, even during maintenance of thalidomide, we think it is unlikely that the manifestation of marrow suppression had been an adverse reaction to the use of these drugs.

The difficulty to define the diagnosis of *A. pullulans* infection, as reported by us and other contemporary authors,^{15,16} owing to agent polymorphism or the large range of differential diagnoses, has confirmed the great value of molecular biology as a diagnostic tool.

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